

MADAP

500 N. Calvert Street, 5th Floor Baltimore, MD 21202 Phone: (410) 767-6535

Fax: (410) 333-2608 or (410) 244-8696

MADAP Prior Authorization Requirements for Hepatitis C Treatment

Purpose: Prior authorization is required for MADAP coverage of Hepatitis C medications to ensure that the patient is eligible for MADAP services and meets the clinical criteria for treatment in accordance with the AASLD/IDSA *HCV Guidance: Recommendations for Testing, Managing, and Treating Hepatitis C.* **Website:** www.hcvguidelines.org

Patient eligibility criteria

- The prescribing clinician must complete the MADAP Prior Authorization Request Form for Hepatitis C Therapy and submit it to MADAP with the applicable documentation to determine the patient's eligibility for MADAP coverage.
- The completed prior authorization request must be faxed to: (410) 333-2608 or (410) 244-8696.
- MADAP must verify that the patient has at least 3 to 6 months of remaining eligibility or is eligible for recertification, depending upon the patient's HCV genotype.
- A patient enrolled in the **Transitional Assistance Program** (**TAP**), while awaiting Medicaid coverage, **is not eligible for MADAP coverage of HCV drugs**.
- MADAP must review the client's insurance coverage, *if applicable*, to determine if the insurance plan is expected to cover some portion of the HCV drug regimen *or* if MADAP is primary.
- As payer of last resort, MADAP will provide coverage for the prescription plan deductibles, co-pays and co-insurance
 for insured clients and the full drug costs for clients who are uninsured or denied coverage by their primary insurance
 plans, within program limits.
- If the patient's Hepatitis C therapy has been approved by MADAP and the patient becomes ineligible for MADAP coverage during therapy, the prescribing clinician must be prepared to enroll the patient in other patient assistance drug programs to complete therapy. (MADAP will make every reasonable effort to maintain coverage until other resources are identified and put into place.)

Clinical and treatment criteria

- The patient must have evidence of chronic hepatitis C infection, with a:
 - Specified genotype and subtype and baseline HCV RNA to determine the course of therapy;
 - Liver biopsy, *FibroSure*[™], *FibroScan*[®] or other comparable HCV test for fibrosis;
 - Prognosis of achieving virologic cure, with treatment, in the judgment of the prescribing clinician.
- The results of the client's liver biopsy, *FibroSure*[™], *FibroScan*® or other comparable test must describe the stage of fibrosis and/or report a Metavir or APRI/FIB4 fibrosis score.
- It is recommended that the patient have an HIV and HCV treatment plan developed and/or medication(s) prescribed in collaboration with a provider who is trained or experienced in treating Hepatitis C or related infectious disease comorbidities, gastroenterology, or hepatology.
 - 1. The patient must be on HIV anti-retroviral therapy ≥ 6 months and/or have an HIV viral load <200 copies/mL within 90 days of having a prior authorization request submitted.
 - 2. The patient should be assessed for potential drug-drug interactions with concomitant medications prior to starting HCV therapy.
 - 3. The HCV RNA viral load should be monitored after 4 weeks of therapy and at 12 weeks following the completion of therapy, per HCV guidelines, to assess the patient's response to the treatment regimen.
- If a ribavirin-containing HCV regimen is prescribed, the patient must utilize 2 forms of contraception while on the regimen and for up to 6 months after stopping, if the patient is:
 - a woman of child-bearing age (at risk for pregnancy), or
 - the male partner of a woman of child-bearing age (at risk for pregnancy).
- The prescribing clinician must certify that an adherence assessment has been performed with the patient and will be conducted throughout the course of treatment to ensure successful completion of the HCV treatment regimen.
- MADAP will pay for one course of treatment, for a current FDA-approved or AASLD recommended drug regimen, not to exceed a 12 to 24-week period, as determined by the patient's HCV genotype, subject to a review of the client's treatment response and/or re-authorization at 6-week intervals.

Please consult the HCV Guidance: Recommendations for Testing, Managing, and Treating Hepatitis C Unique Patient Populations: Patients with HIV/HCV Coinfection

http://www.hcvguidelines.org/full-report/unique-patient-populations-patients-hivhcv-coinfection



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	Client Services Use Only	
Date Rec'd:	Insurance Review:	
Date Rev'd:	□ Approved: □ Denied:	
Elig. Date: PA Dates:		
PA entered:/	/ PA#: Int:	

Fax completed form to: (410) 333-2608 or (410) 244-8696

MADAP PRIOR AUTHORIZATION REQUEST FORM FOR HEPATITIS C THERAPY

Part 1: Patient Information

Name:	MADAP Client ID #: 9 4					
DOB:/Body Weight:	kg Pt's Daytime Phone #:					
What is the expected or actual start date of the patient's Hepatitis C treatment regimen?/						
Part 2: Hepatitis C Treatment Plan						
Check the requested treatment regimen from the following selections.						
☐ Daklinza ® (daclatasvir) mg: Take once daily for weeks	☐ Harvoni ® (ledipasvir-sofosbuvir) 90 mg/400 mg: Take tablet(s) once daily for weeks					
☐ Olysio ® (simeprevir) 150 mg: Take once daily for weeks	☐ Sovaldi ® (sofosbuvir) 400 mg: Take once daily for weeks					
☐ Technivie ® (ombitasvir/paritaprevir/ritonavir) 25 mg/150 mg/ 100 mg: Take tablet(s) once daily for weeks	☐ Viekira Pak® (dasabuvir) 250 mg + (ombitasvir/paritaprevir/ritonavir) 25 mg/150 mg/100 mg: Take as directed for weeks					
☐ Zepatier ® (elbasvir/grazoprevir) 50 mg/100 mg: Take once daily for weeks	☐ Peginterferon alfa mcg: Inject once weekly for weeks					
□ Ribavirin mg: Take in the	□: eks Take: weeks					
morning and in the afternoon for we						
☐ as directed for weeks	:					
Part 3: Hepatitis C Diagnostic Information Complete <u>all</u> of the diagnostic information that follows as it applies to this patient.						
What is the patient's HCV genotype and subtype?						
Baseline quantitative HCV RNA (within 90 days of request date): IU/mL; Test date://						
Has a fibrosis test been performed? ☐ No ☐	Yes; Test date:/; Test used:					
If Yes, Metavir Grade:; Metavir Stag	ge:, if applicable					
Has a liver biopsy been performed? ☐ No ☐	☐ Yes; Test date:/					
If Yes, Metavir Grade:; Metavir Stag	ge:					
\Rightarrow A copy of the HCV genotype, RNA, fibrosis and/or biopsy test results must be provided with this request. \Rightarrow (If performed, please include a copy of the patient's test results for the presence of resistance-associated variants.)						
	in the judgment of the prescribing clinician has a prognosis of Yes, select one of the following:					
☐ Acute Hepatitis C infection	☐ Liver transplant recipient:					
(HCV seroconversion or discrete exposure less than 6 months prior to request date)	HCV genotype and subtype of pre-transplant liver:					
☐ Chronic Hepatitis C	HCV genotype and subtype of post-transplant liver:					
☐ Hepatocellular Carcinoma	☐ Other:					

Part 4: Hepatitis C Treatment History

Has this patient been treated for Hepatitis C If Treatment Experienced, what we	in the past? Treatment Naïvovas the outcome of the previous treatment		ment Experienced		
☐ Relapsed ☐ Partial Re Please indicate what prior regimen(s) the partial Re	sponder	☐ Toxic	ities		
HCV regimen	Treatment duration/ dates		ment Outcome		
		☐ Relapsed☐ Non-Responder☐ Other:	□ Partial Responder □ Toxicities		
		□ Relapsed □ Non-Responder □ Other:			
	Part 5: HIV Treatment Statu	s			
HIV RNA viral load (within 90 days of requ	uest date):copies/ml	Date://	_		
Has the patient been on HIV anti-retroviral therapy ≥ 6 months? \square No \square Yes					
Patient is expected to take HIV medications during HCV treatment:					
If No, state reason:					
If Yes, list the ART being prescribed:					
Part 6: Precautions Review and Adherence Assessment					
Has a pre-treatment precautions review and Patient has an active diagnosis of a	med? □ No □ No	☐ Yes, <i>please specify:</i> ☐ Yes			
Patient is engaged in or has been re	t. 🗆 No	☐ Yes ☐ NA			
Pregnancy-related precautions wer	□ No	☐ Yes ☐ NA			
Will precautions and adherence monitoring continue during the course of HCV treatment? ☐ No ☐ Yes					
Please describe any other precautions and/or adherence issues that were reviewed with the patient:					
Part 7: Patient Assistance Affidavit					
Is the prescribing clinician prepared to have the patient enrolled in other patient assistant drug programs to complete the Hepatitis C therapy, if MADAP has approved the HCV regimen, but the patient subsequently becomes ineligible for MADAP coverage?					
I certify that the information provided is accurate. Supporting documentation is available for State audits.					
Prescribing Clinician:					
Clinic/Hospital Name:					
Street Address:					
City, State & Zip:					
Clinician's Signature:		Date: _	//		